AMENDMENTS TO THE CLAIMS

1-15. (Cancelled)

16. (Currently Amended) A method for infusing a fluid in a living body, said method comprising:

providing a reservoir, a flow restrictor, and a valve in an implantable drug pump device:

transiently storing [[a]] fluid infusate in said reservoir for transmission to a delivery site after the implantable drug pump device has been implanted in a patient;

limiting a flow rate of the fluid infusate using said flow restrictor disposed in a fluid path between said reservoir and said delivery site;

determining transient pressure differentials relative to said flow restrictor by a controller component of the implantable drug pump;

determining whether an occlusion is present in the flow path using the transient pressure differentials, the transient pressure differentials being determined when the controller component closes a valve to interrupt flow of the fluid infusate from the implantable drug pump device; and

determining a location of an occlusion when an occlusion is detected using the transient pressure differentials;

controlling said valve disposed in said fluid path between said reservoir and said delivery site to control infusate output from said reservoir to said delivery site as a function of said transient pressure differentials across said flow restrictor, wherein the controlling said valve automatically responds to a detection of an a partial occlusion by altering a unit dose period for delivery of the fluid infusate;

communicating a signal, from implantable drug pump device to an external device, that is indicative of an amount of occlusion detected by the controller; and

communicating a signal, from implantable drug pump device to an external device, that is indicative of a location of the occlusion.

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17. (Previously Presented) The method of claim 16, wherein controlling said valve comprises:

subdividing a flow period into smaller unit dose periods over which a pressure differential across said flow restrictor is likely to remain constant and controlling said valve to deliver a total dose of said infusate through a series of sequential said unit dose periods.

- 18. (Original) The method of claim 17, wherein said unit dose periods are selected at least in part to reduce battery consumption.
- 19. (Original) The method of claim 17, wherein said unit dose periods are selected so that an open/close rate of said valve is pharmacologically insignificant.
- 20. (Previously Presented) The method of claim 16, further comprising: providing an alert with respect to overfilling of said reservoir using a pressure differential across said flow restrictor.
- 21. (Previously Presented) The method of claim 16, further comprising: providing an alert with respect to depletion of said reservoir using a pressure differential across said flow restrictor.
 - 22. (Previously Presented) The method of claim 16, further comprising: determining a rate at which a pressure differential across said flow restrictor changes.
- 23. (Previously Presented) The method of claim 22, wherein controlling said valve comprises:

altering timing of a period of said valve being opened as a function of said rate at which a pressure differential across said flow restrictor changes.

24. (Original) The method of claim 16, further comprising: determining a temperature of said infusate, wherein controlling said valve comprises controlling infusate output from said reservoir as a function of said temperature of said infusate. Application No.: 10/626,902 Docket No. 02-029 CIP

25-34. (Cancelled)